Food and Drug Administration, HHS

§26.81 Final provisions.

- (a) The sectoral annexes referred to in §26.80(a), as well as any new sectoral annexes added pursuant to §26.80(b), shall form an integral part of the "Agreement on Mutual Recognition Between the United States of America and the European Community," from which this part is derived.
- (b) For a given product or sector, the provisions contained in subparts A and B of this part shall apply in the first place, and the provisions of subpart C of this part in addition to those provisions. In the case of any inconsistency between the provisions of subpart A or B of this part and subpart C of this part, subpart A or B shall prevail, to the extent of that inconsistency.
- (c) The agreement from which this part is derived shall not affect the rights and obligations of the parties under any other international agreement.
- (d) In the case of subpart B of this part, the parties shall review the status of such subpart at the end of 3 years from the date described in §26.80(a).

PART 50—PROTECTION OF HUMAN SUBJECTS

Subpart A—General Provisions

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- 50.55 Requirements for permission by parents or guardians and for assent by children.

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AUTHORITY: 21 U.S.C 321, 343, 346, 346a, 348, 350a, 350b, 352, 353, 355, 360, 360c-360f, 360h-360j, 371, 379e, 381; 42 U.S.C. 216, 241, 262, 263b-263n.

Source: 45 FR 36390, May 30, 1980, unless otherwise noted.

Subpart A—General Provisions

§ 50.1 Scope.

(a) This part applies to all clinical investigations regulated by the Food and Drug Administration under sections 505(i) and 520(g) of the Federal Food, Drug, and Cosmetic Act, as well as clinical investigations that support applications for research or marketing permits for products regulated by the Food and Drug Administration, including foods, including dietary supplements, that bear a nutrient content claim or a health claim, infant formulas, food and color additives, drugs for human use, medical devices for human use, biological products for human use, and electronic products. Additional specific obligations and commitments of, and standards of conduct for, persons who sponsor or monitor clinical investigations involving particular test articles may also be found in other parts (e.g., parts 312 and 812). Compliance with these parts is intended to protect the rights and safety of subjects involved in investigations filed with the Food and Drug Administration pursuant to sections 403, 406, 409, 412, 413, 502, 503, 505, 510, 513-516, 518-520, 721, and 801 of the Federal Food, Drug, and Cosmetic Act and sections 351 and 354-360F of the Public Health Service Act.

(b) References in this part to regulatory sections of the Code of Federal